

# Coiled-coil structure-dependent interactions between polyQ proteins and Foxo lead to dendrite pathology and behavioral defects

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Neurodegenerative disorders, such as Huntington's diseases and spinocerebellar ataxias (SCAs), are driven by proteins with expanded polyglutamine (polyQ) tracts. Recently, coiled-coil structures in polyQ regions of such proteins were shown to facilitate aggregate formation and ultimately lead to cell death. However, the molecular mechanism linking these structural domains to neuronal toxicity of polyQ proteins remains elusive. Here, we demonstrate that coiled-coil structures in the Q repeat region of SCA type 3 (SCA3) polyQ proteins confer protein toxicity in *Drosophila* neurons. To functionally characterize coiled-coil structures in the Q repeat regions, we generated three structural variants of SCA3 polyQ proteins: (i) MJDtr-76Q, containing both  $\alpha$ -helical coiled-coil and  $\beta$ -sheet hairpin structures in the Q repeat region; (ii) MJDtr-70Q<sub>cc0</sub>, possessing only  $\alpha$ -helical coiled-coil structures due to the incorporation of  $\beta$ -sheet-breaking residues (Q-to-N or Q-to-E mutations); and (iii) MJDtr-70Q<sub>pqp</sub>, with no secondary structure due to the introduced proline residues (Q-to-P mutations). Through comparative analysis of these variants, we found that coiled-coil structures facilitated nuclear localization of SCA3 polyQ proteins and induced dendrite defects in *Drosophila* dendritic arborization neurons. Furthermore, genetic and functional screening identified the transcription factor Foxo as a target of polyQ proteins, and coiled-coil-mediated interactions of Foxo and polyQ proteins in the nucleus resulted in the observed dendrite and behavioral defects in *Drosophila*. These results demonstrate that coiled-coil structures of polyQ proteins are crucial for their neuronal toxicity, which is conferred through coiled-coil to coiled-coil interactions with the nuclear targets of these proteins.

polyQ | coiled coil | Foxo | dendrites | neurodegenerative diseases

**P**olyglutamine (polyQ) diseases are neurodegenerative disorders that are characterized by the expansion of Q repeats in the associated proteins, such as those underlying spinocerebellar ataxias (SCAs) and the Huntingtin protein (HTT) (1, 2). The accumulation of misfolded polyQ proteins caused by Q expansion confers protein toxicity that results in neuropathic features, such as dendrite defects, leading to impaired neuronal functions and neuronal cell death (3, 4). Numerous studies (5, 6) have reported that aberrant dendritic morphologies are linked to defects in brain function, such as memory storage and cognition. Because pathological dendritic features are often observed in patients with polyQ diseases (7, 8), it is important to understand how misfolded polyQ proteins contribute to dendrite pathology.

Cellular and molecular mechanisms linking polyQ protein toxicity to dendrite pathology have been uncovered in models of several polyQ diseases. PolyQ protein toxicity can be conferred through the impairment of several cellular components, such as

the endosomal machinery or cytoskeleton. For example, Richards et al. (9) showed that dendrite defects were associated with perturbations of the Rab11-dependent endosomal recycling system in a *Drosophila* model of Huntington's disease. In addition, we previously demonstrated that dendrite defects were associated with perturbed actin cytoskeletal structure and impaired subcellular distribution of Golgi outposts in *Drosophila* models of SCA type 3 (SCA3), also known as Machado-Joseph disease (MJD) (10, 11). Despite these efforts, the molecular mechanism of polyQ protein toxicity and the link to the resulting dendrite pathology remain largely unknown.

The structure of polyQ proteins is one of the determining factors in the associated toxicity, but it is difficult to uncover the structural features of misfolded polyQ proteins using X-ray

## Significance

It remains unclear how the structural properties of polyglutamine (polyQ) proteins, which underlie several neurodegenerative disorders, including Huntington's disease and spinocerebellar ataxias (SCAs), translate into the toxicity of these proteins. Here, we demonstrate that coiled-coil structures in expanded polyQ regions of SCA type 3 (SCA3) proteins cause dendrite defects in *Drosophila* neurons, as well as behavioral abnormalities. Moreover, interactions of SCA3 with Foxo mediated by coiled-coil domains of these two proteins resulted in functional impairment of this transcription factor, whereas its overexpression significantly rescued the SCA3-induced defects. Our study expanded the current understanding of neuronal pathology mediated by polyQ proteins via the coiled-coil-mediated interactions. These results may have important implications in therapeutic strategies for polyQ protein-related diseases.

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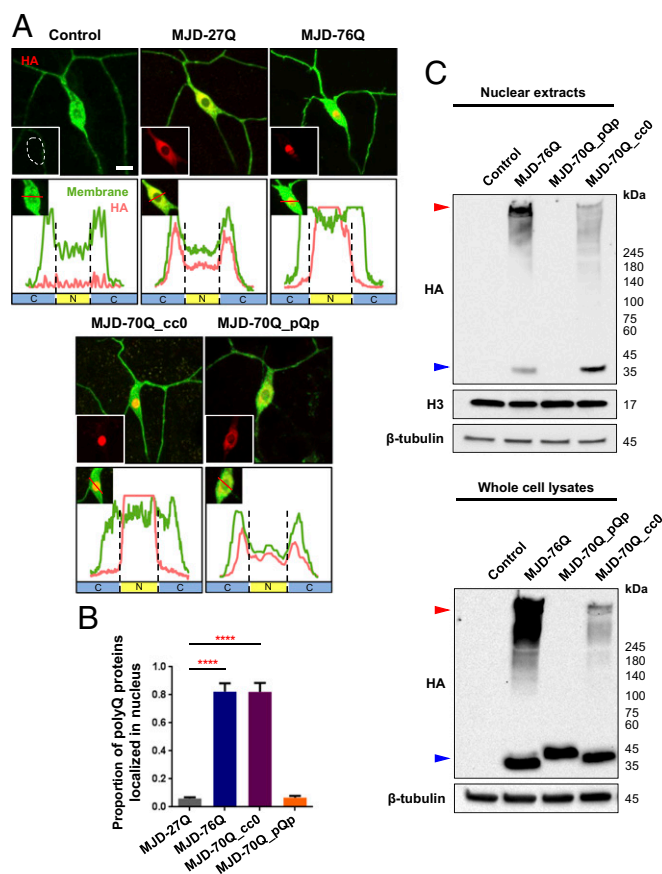




cytoskeletal structures in da neuronal clusters, and found that F-actin cytoskeletal structures of distal dendrites were substantially disrupted in da neuronal clusters expressing MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> but not in those expressing MJDtr-70Q<sub>pQp</sub>, compared with those in the control da neuronal clusters (*SI Appendix, Fig. S1 G and H*). In another previous study, we showed that the plasma membrane (PM) protein supply was severely reduced by pathogenic SCA3 polyQ proteins, leading to impaired elongation of dendrite terminals in C4da neurons (11). We thus examined alterations of the PM protein supply in C4da neurons expressing the three structural variants and found that this supply was diminished in the neurons expressing MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> but not in those expressing MJDtr-70Q<sub>pQp</sub> (Fig. 1E and *SI Appendix, Fig. S1I*). Taken together, these results suggest that coiled-coil structures in the Q repeat regions of MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins contribute to pathological dendrite phenotypes, including aberrant terminal dendrite morphology, impaired F-actin cytoskeleton, and diminished PM protein supply.

**Coiled-Coil Structures of SCA3 polyQ Proteins Promote Their Nuclear Localization.** Nuclear localization of SCA3 polyQ proteins is critical for their toxicity and the associated dendrite defects in C4da neurons (10, 11). As the coiled-coil structures contributed to this toxicity, we next asked whether coiled-coil structures might facilitate the localization of SCA3 polyQ proteins to the nucleus. To this end, we examined the subcellular localization of the three SCA3 structural variants in C4da neurons, and found that MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins were primarily localized to the nucleus, whereas MJDtr-70Q<sub>pQp</sub> proteins were diffusely distributed in the cytoplasm (Fig. 2A). The nuclear-localized proportions of MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins were significantly ( $P < 1.0 \times 10^{-4}$ ) larger than those of MJDtr-70Q<sub>pQp</sub> proteins (Fig. 2B). We also examined the subcellular localization of MJDtr-27Q proteins, as a nonpathogenic control, that have coiled-coil domains with a normal repeat of Qs (*SI Appendix, Fig. S2A*) (18). MJDtr-27Q proteins were diffusely distributed in the cytoplasm with a marginal nuclear-localized proportion, similar to MJDtr-70Q<sub>pQp</sub> proteins (Fig. 2A and B). These results suggest that coiled-coil domains with only expanded repeats of Qs (MJDtr-76Q and MJDtr-70Q<sub>cc0</sub>) appear to facilitate the nuclear localization of SCA3 polyQ proteins, although other biophysical properties of polyQ proteins may also contribute to their subcellular localization. To corroborate this finding, we performed Western blot analysis of nuclear lysates from pan-neuronal cells expressing the three SCA3 structural variants using the *elav-Gal4* driver. Consistently, significant amounts of MJDtr-70Q<sub>cc0</sub> and MJDtr-76Q proteins, but not MJDtr-70Q<sub>pQp</sub> proteins, were detected in the nuclear lysates (Fig. 2C). MJDtr-70Q<sub>pQp</sub> proteins were detected only in cytoplasmic lysates (*SI Appendix, Fig. S2B*). Certain amounts of monomers and insoluble aggregates of MJDtr-70Q<sub>cc0</sub> and MJDtr-76Q were detected in cytoplasmic lysates, suggesting that these two proteins are primarily enriched in the nucleus, but are not absent from the cytoplasm, as also indicated by the immunohistochemistry data (Fig. 2A).

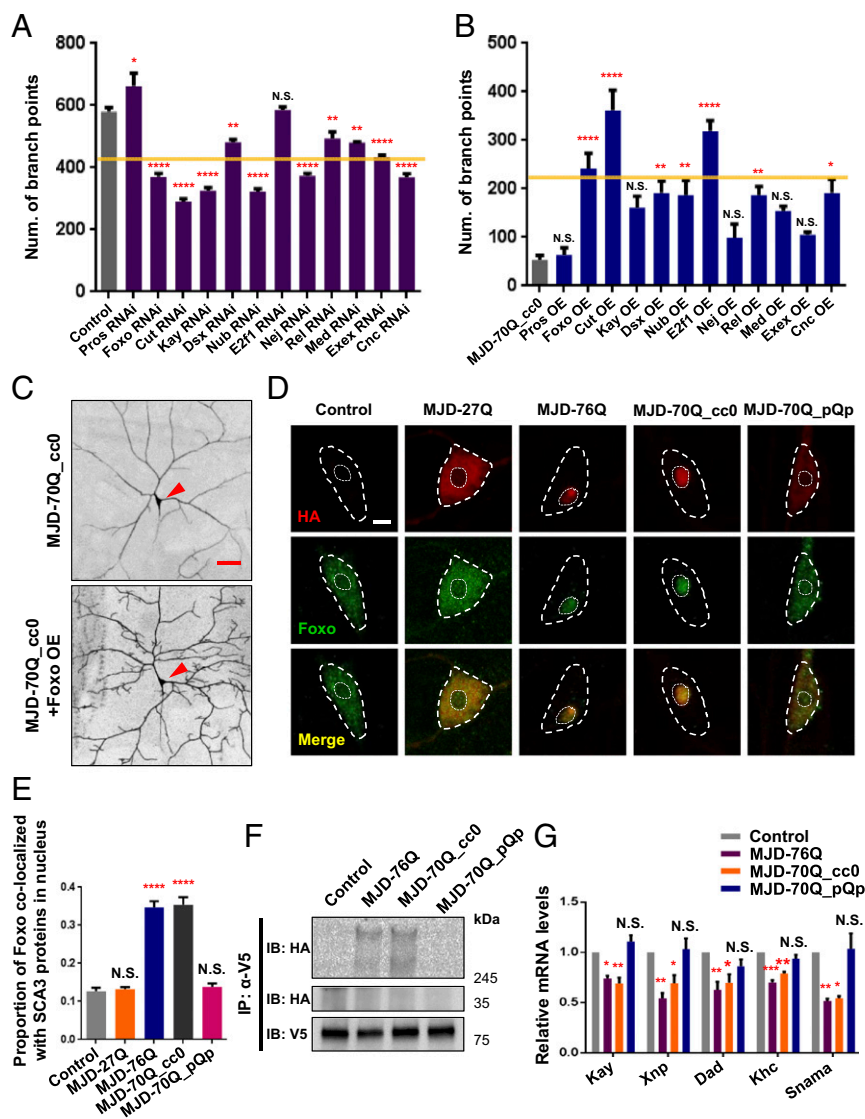
**Interaction of Forkhead Box, Subgroup O with SCA3 polyQ Proteins in the Nucleus Leads to Its Functional Impairment and Ensuing Dendrite Defects.** Next, we searched for molecular targets of nuclear-localized MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins that mediated the observed dendrite defects. It has been suggested that polyQ proteins can target cellular proteins through coiled-coil to coiled-coil interactions (19). Furthermore, it was previously reported that the targets of mutant HTT proteins had a tendency to contain high amounts of coiled-coil structures (14). Based on these reports, we hypothesized that coiled-coil structures in the Q repeat region mediate interactions of nuclear MJDtr-76Q and



**Fig. 2.** Coiled-coil structures of SCA3 polyQ proteins promote their nuclear localization. (A) Immunohistochemical images representing subcellular localization of SCA3 proteins in C4da neurons expressing MJDtr-27Q, MJDtr-76Q, MJDtr-70Q<sub>pQp</sub>, or MJDtr-70Q<sub>cc0</sub> proteins (*UAS-CD4td-GFP/+;ppk-Gal4/+; UAS-CD4td-GFP/+;ppk-Gal4/UAS-MJDtr-27Q, UAS-CD4td-GFP/+;ppk-Gal4/UAS-MJDtr-76Q, UAS-CD4td-GFP/+;ppk-Gal4/UAS-MJDtr-70Q<sub>cc0</sub>, and UAS-CD4td-GFP/+;ppk-Gal4/UAS-MJDtr-70Q<sub>pQp</sub>*). (Upper five panels of each denoted genotype) Merged images of SCA3 proteins (red) with membrane marker, mCD8-GFP (green). Left Lower Insets represent cell body regions, and the white dashed line in the control C4da neuron indicates the outline of a cell body. (Scale bar, 5  $\mu$ m.) (Lower five panels of each denoted genotype) Intensity profiles of fluorescent signals representing SCA3 proteins (red) and mCD8-GFP (green) across cell bodies along red lines (Left Upper Insets). Black dashed lines distinguish nucleus and cytoplasm. N (yellow) indicates nucleus. C (blue) indicates cytoplasm. (B) Comparison of nuclear proportions of SCA3 polyQ proteins among C4da neurons expressing transgenes described in A. \*\*\*\* $P < 1.0 \times 10^{-4}$  by one-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 6$  neurons. (C) Western blot analyses using whole cell lysates and nuclear extracts of adult fly heads (*elav-Gal4/+; elav-Gal4/MJDtr-76Q, elav-Gal4/MJDtr-70Q<sub>cc0</sub>, and elav-Gal4/MJDtr-70Q<sub>pQp</sub>*). SCA3 variants were immunoblotted using anti-HA antibody. Histone 3 (H3) is used as the loading control for nuclear extracts, and  $\beta$ -tubulin is used as the loading controls for whole cell lysates and nuclear extracts. Red arrowheads indicate insoluble aggregates of SCA3 polyQ proteins. Blue arrowheads indicate monomers of SCA3 polyQ proteins.  $n = 3$  independent experiments.

MJDtr-70Q<sub>cc0</sub> proteins with other coiled-coil-rich proteins. Thus, we searched for transcription factors (TFs) that could regulate dendrite morphology and interact with MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins. To this end, we first identified 58 TFs that are involved in dendrite morphogenesis based on gene ontology biological processes and the previous literature (11, 20–26). We then selected 13 TFs with the largest numbers of amino acids in their coiled-coil domains (Table 1), and these TFs were tested for their effect on dendrite morphology. Among the 13 TFs, knockdown of 6 TFs [Forkhead box, subgroup O (Foxo),



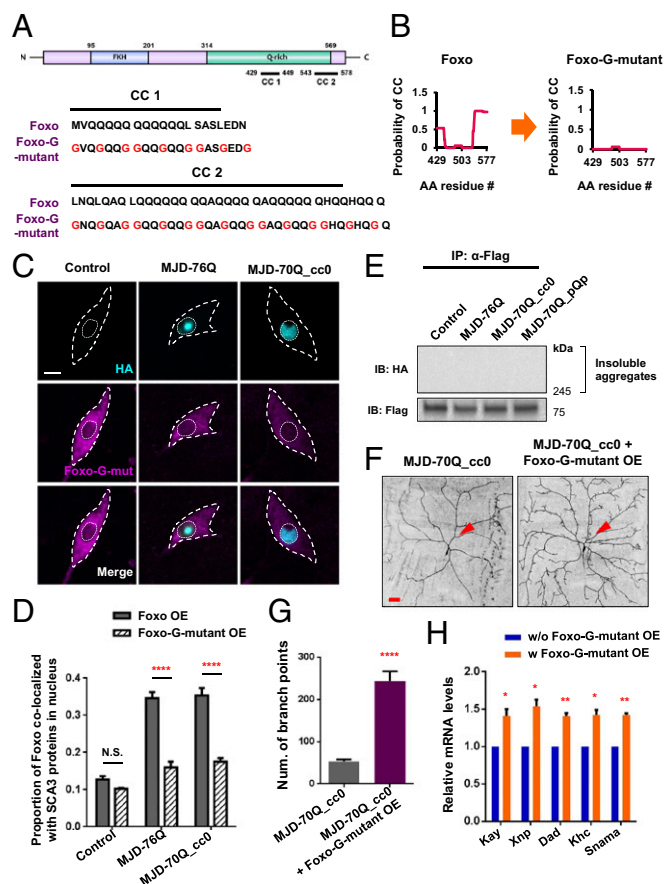


**Fig. 3.** Coiled-coil-rich Foxo interacts with SCA3 polyQ proteins, leading to its functional impairment. (A) Quantification of the number of dendrite branch points in C4da neurons expressing the denoted RNAi lines (see *Materials and Methods*). Each RNAi and *UAS-CD4-tdGFP* were expressed using *ppk1<sup>ts</sup>-Gal4*. The yellow line indicates 25% reduction in the number of dendrite branch points compared with that in control C4da neurons. N.S., not significant; \* $P < 0.05$ ; \*\* $P < 1.0 \times 10^{-2}$ ; \*\*\*\* $P < 1.0 \times 10^{-4}$  by one-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 3$  neurons. (B) Quantification of the number of dendrite branch points in C4da neurons overexpressing MJDtr-70Q\_cc0 or MJD-70Q\_cc0+ the denoted TFs (see *Materials and Methods*). Each TF and *UAS-MJDtr-70Q\_cc0* was coexpressed by *ppk-Gal4*. The yellow line indicates a fourfold increase in the number of dendrite branch points compared with that in C4da neurons expressing MJDtr-70Q\_cc0 alone. N.S., not significant; \* $P < 0.05$ ; \*\* $P < 1.0 \times 10^{-2}$ ; \*\*\*\* $P < 1.0 \times 10^{-4}$  by one-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 3$  neurons. (C) Dendrite images of C4da neurons expressing MJDtr-70Q\_cc0 (Upper) or MJDtr-70Q\_cc0 + Foxo (Lower) (*UAS-CD4-tdGFP/+; ppk-Gal4,UAS-MJDtr-70Q\_cc0/+* and *UAS-CD4-tdGFP/UAS-Foxo;ppk-Gal4,UAS-MJDtr-70Q\_cc0/+*). Arrowheads (red) indicate cell bodies. (Scale bar, 50  $\mu$ m.) (D) Subcellular localization of Foxo (green) and SCA3 structural variants (red) in C4da neurons overexpressing Foxo, MJDtr-27Q + Foxo, MJDtr-76Q + Foxo, MJDtr-70Q\_cc0 + Foxo, or MJDtr-70Q\_pQp + Foxo (*ppk-Gal4,UASp-Foxo-GFP/+*, *ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-27Q*, *ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-76Q*, *ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-70Q\_cc0*, and *ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-70Q\_pQp*). (Scale bar, 5  $\mu$ m.) Outer and inner dashed lines (white) indicate the outlines of the cell body and nucleus. (E) Comparison of nuclear proportions of Foxo among C4da neurons expressing transgenes described in D. N.S., not significant; \*\*\*\* $P < 1.0 \times 10^{-4}$  by one-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 12$  neurons. (F) Representative image of coimmunoprecipitation experiments using adult fly heads (*UAS-Foxo-V5/+;elav-GeneSwitch-Gal4/+*, *UAS-Foxo-V5/+;elav-GeneSwitch-Gal4/UAS-MJDtr-76Q*, *UAS-Foxo-V5/+;elav-GeneSwitch-Gal4/UAS-MJDtr-70Q\_cc0*, and *UAS-Foxo-V5/+;elav-GeneSwitch-Gal4/UAS-MJDtr-70Q\_pQp*). Top and Middle indicate immunoprecipitated insoluble aggregates and monomeric forms of HA-tagged SCA3 proteins, respectively. Bottom shows V5-tagged Foxo proteins.  $n = 3$  independent experiments. (G) mRNA expression levels of Foxo target genes (*Kay*, *Xnp*, *Dad*, *Khc*, and *Snama*) measured by RT-PCR analysis in adult fly heads (*elav-Gal4/+*, *elav-Gal4/UAS-MJDtr-76Q*, *elav-Gal4/UAS-MJDtr-70Q\_cc0*, and *elav-Gal4/UAS-MJDtr-70Q\_pQp*). N.S., not significant; \* $P < 0.05$ ; \*\* $P < 1.0 \times 10^{-2}$ ; \*\*\*\* $P < 1.0 \times 10^{-3}$  by two-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 3$  independent experiments.

whether Foxo-G mutants could restore dendrite defects caused by MJDtr-70Q\_cc0 proteins. Overexpression of Foxo-G mutant proteins in C4da neurons expressing MJDtr-70Q\_cc0 significantly ( $P < 1.0 \times 10^{-4}$ ) restored the defects in dendrite morphology and the number of dendrite branch points (Fig. 4 F

and G). We then examined whether the transcriptional activity of Foxo reduced by MJDtr-70Q\_cc0 proteins could also be restored by overexpression of Foxo-G mutant proteins. mRNA levels of Foxo target genes (*Kay*, *Xnp*, *Dad*, *Khc*, and *Snama*) down-regulated by MJDtr-70Q\_cc0 proteins were significantly ( $P < 0.05$ ) restored by





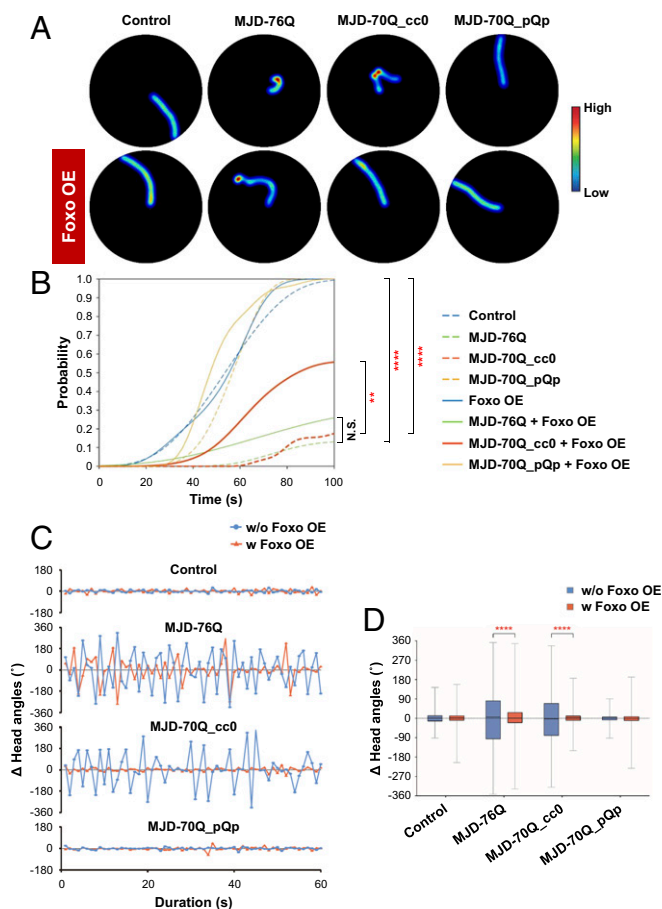
**Fig. 4.** Disruption of coiled-coil structures in Foxo abolishes its interaction with SCA3 polyQ proteins. (A) Amino acid changes in Foxo-G mutants within the Q-rich domain of Foxo. Foxo contains a Forkhead (FKH) domain—a DNA-binding domain—in the N-terminal region, and a Q-rich domain including two coiled-coil domains (CC1 and CC2) of wild-type Foxo and Foxo-G mutant proteins are shown in a heptad repeat pattern. *a/d/g* residues of coiled-coil heptad repeats are replaced with glycines (Gs), which are known as coiled-coil breakers. Replaced amino acids are marked in red. (B) Probabilities to form coiled-coil structure in Foxo-G mutants calculated by COILS. The y axis indicates probability to form coiled-coil structure, and x axis indicates the amino acid residue numbers. (C) Subcellular localization of Foxo-G mutant proteins (magenta) and SCA3 structural variants (cyan) in C4da neurons overexpressing Foxo-G mutant, MJDtr-76Q + Foxo-G mutant, or MJDtr-70Q<sub>cc0</sub> + Foxo-G mutant (*UAS-Foxo-G mutant-Flag/UAS-CD4-tGFP;ppk-Gal4/+*, *UAS-Foxo-G mutant-Flag/UAS-CD4-tGFP;ppk-Gal4,UAS-MJDtr-76Q/+*, and *UAS-Foxo-G mutant-Flag/UAS-CD4-tGFP;ppk-Gal4,UAS-MJDtr-70Q<sub>cc0</sub>/+*). (Scale bar, 5  $\mu$ m.) Outer and inner dashed lines (white) indicate the outlines of the cell body and nucleus. (D) Comparison of nuclear proportions of wild-type Foxo and Foxo-G mutants colocalized with SCA3 proteins among C4da neurons expressing the transgenes described in C. N.S., not significant; \*\*\*\* $P < 1.0 \times 10^{-4}$  by two-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 12$  neurons. (E) Representative image of co-IP experiments using adult fly heads (*UAS-Foxo-G mutant-Flag/+*; *elav-GeneSwitch-Gal4/+*, *UAS-Foxo-G mutant-Flag/+*; *elav-GeneSwitch-Gal4/UAS-MJDtr-76Q*, *UAS-Foxo-G mutant-Flag/+*; *elav-GeneSwitch-Gal4/UAS-MJDtr-70Q<sub>cc0</sub>*, and *UAS-Foxo-G mutant-Flag/+*; *elav-GeneSwitch-Gal4/UAS-MJDtr-70Q<sub>pQp</sub>*). Upper and Lower panels show insoluble aggregates of HA-tagged SCA3 proteins and Flag-tagged Foxo-G mutant proteins, respectively.  $n = 3$  independent experiments. (F) Dendrite images of C4da neurons expressing MJDtr-70Q<sub>cc0</sub> (Left) or MJDtr-70Q<sub>cc0</sub> + Foxo-G mutant (Right) (*UAS-CD4-tGFP/+*; *ppk-Gal4,UAS-MJDtr-70Q<sub>cc0</sub>/+* and *UAS-CD4-tGFP/+*; *ppk-Gal4,UAS-MJDtr-70Q<sub>cc0</sub>/+*; *UAS-CD4-tGFP/+*; *ppk-Gal4,UAS-MJDtr-70Q<sub>cc0</sub>/+*; *UAS-CD4-tGFP/+*; *ppk-Gal4,UAS-MJDtr-70Q<sub>cc0</sub>/+*; *UAS-CD4-tGFP/+*; *ppk-Gal4,UAS-MJDtr-70Q<sub>cc0</sub>/+*). Arrowheads (red) indicate cell bodies. (Scale bar, 50  $\mu$ m.) (G) Quantification of the number of dendrite branch points of C4da neurons expressing the transgenes described in F. \*\*\*\* $P < 1.0 \times 10^{-4}$  by Student's *t* test; error bars, SEM;  $n \geq 5$  neurons. (H) mRNA levels of Foxo target genes (*Kay*, *Xnp*, *Dad*, *Khc*, *Snama*) measured by RT-PCR analysis in adult fly heads (*elav-Gal4/UAS-MJDtr-70Q<sub>cc0</sub>* and *UAS-Foxo-G mutant-Flag/+*; *elav-Gal4/UAS-MJDtr-70Q<sub>cc0</sub>*). \* $P < 0.05$ ; \*\* $P < 1.0 \times 10^{-2}$  by Student's *t* test; error bars, SEM;  $n = 3$  independent experiments.

co-overexpression of Foxo-G mutant proteins (Fig. 4H). These data indicate that coiled-coil to coiled-coil interactions mediate the interaction of Foxo with MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins, contributing to dendrite pathology.

**Foxo Overexpression Restores the Behavioral Defects Induced by Coiled-Coil Structure-Mediated polyQ Protein Toxicity.** In addition to causing dendrite defects (Fig. 1C), polyQ proteins have been shown to affect neuronal survival, animal motility, and animal viability (3, 27). Thus, we investigated whether Foxo could rescue the motility defects induced by polyQ proteins in *Drosophila* larvae. Larval motility is assessed by measuring crawling and turning activities (28). We first tested the crawling capability using the following eight different lines of larvae: control larvae, and larvae expressing MJDtr-76Q, MJDtr-70Q<sub>cc0</sub>, or MJDtr-70Q<sub>pQp</sub> proteins in da neuronal clusters with or without co-overexpression of Foxo using the *109(2)80-gal4* driver. For each line, we placed a larva in the center of a Petri dish and monitored its movements until it reached the edge of the dish for up to 100 s (Fig. 5A) and repeated this experiment with more than 20 different larvae. We measured the time for the larvae to reach the dish edge and the cumulative fraction of larvae reaching the dish edge over time. All control larvae reached the dish edge within 100 s (Fig. 5B). However, only 13.3% and 19.2% of larvae expressing MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins, respectively, reached the dish edge within 100 s (Fig. 5B). The cumulative fraction of larvae co-overexpressing Foxo and MJDtr-70Q<sub>cc0</sub> proteins significantly ( $P < 1.0 \times 10^{-4}$ ) increased to 56.3%, while larvae overexpressing Foxo alone showed no significant difference from the control, suggesting that Foxo is important in the regulation of larval motility affected by SCA3 polyQ proteins. In contrast, the cumulative fraction of larvae expressing MJDtr-70Q<sub>pQp</sub> proteins was not significantly different from that of control larvae (Fig. 5B), suggesting that the crawling defects were induced by coiled-coil structures in polyQ proteins. Co-overexpression of Foxo and MJDtr-76Q resulted in only a partial (2.28-fold) increase in the cumulative fraction compared with larvae overexpressing MJDtr-76Q alone, although this increase was not statistically significant (Fig. 5B). This partial restoration suggests a potential additional toxicity mediated by coiled-coil structure-independent properties of polyQ proteins.

We next used the same transgenic lines to examine another behavior: the larval head turning. To analyze the characteristics of head turning, we monitored the changes in the angle of larval heads for 1 min (*SI Appendix, Fig. S5A*). Control larvae rarely turned their heads, but larvae expressing MJDtr-70Q<sub>cc0</sub> showed significantly ( $P < 1.0 \times 10^{-4}$ ) more frequent turning of their heads, while larvae expressing MJDtr-70Q<sub>pQp</sub> showed patterns of head turning similar to those of control larvae (Fig. 5C and D). The defects in head turning were significantly ( $P < 1.0 \times 10^{-4}$ ) restored by Foxo overexpression in larvae expressing MJDtr-76Q or MJDtr-70Q<sub>cc0</sub>, but the effect was stronger in larvae expressing MJDtr-70Q<sub>cc0</sub> (Fig. 5C and D). The overexpression of Foxo alone had no effect on head turning (*Discussion* and Fig. 5). We next examined whether loss-of-function of Foxo could lead to behavioral defects similar to those induced by MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins. Indeed, larvae expressing Foxo RNAi showed similar defects in both crawling and turning (*SI Appendix, Fig. S5B–E*). Taken together, these data suggest that Foxo can rescue the defects induced by coiled-coil structure-mediated polyQ

*Dad*, *Khc*, and *Snama*) measured by RT-PCR analysis in adult fly heads (*elav-Gal4/UAS-MJDtr-70Q<sub>cc0</sub>* and *UAS-Foxo-G mutant-Flag/+*; *elav-Gal4/UAS-MJDtr-70Q<sub>cc0</sub>*). \* $P < 0.05$ ; \*\* $P < 1.0 \times 10^{-2}$  by Student's *t* test; error bars, SEM;  $n = 3$  independent experiments.



**Fig. 5.** Foxo overexpression restores larval locomotion defects induced by coiled-coil structure-mediated polyQ toxicity. (A) Heat maps showing residence probability during traveling of larvae [109(2)80-Gal4/+; +/+; 109(2)80-Gal4/+; UAS-MJDtr-76Q/+; 109(2)80-Gal4/+; UAS-MJDtr-70Q<sub>cc0</sub>/+, 109(2)80-Gal4/+; UAS-MJDtr-70Q<sub>pQp</sub>/+, 109(2)80-Gal4/+; UAS-Foxo/+; 109(2)80-Gal4/+; UAS-Foxo/+; UAS-MJDtr-76Q/+; 109(2)80-Gal4/+; UAS-Foxo/+; UAS-MJDtr-70Q<sub>cc0</sub>/+, and 109(2)80-Gal4/+; UAS-Foxo/+; UAS-MJDtr-70Q<sub>pQp</sub>/+] monitored in the 90-mm Petri dish for up to 100 s. (B) Cumulative density functions showing fractions of larvae that reached the dish edge over time. The functions are estimated separately for larvae expressing each of the transgenes described in A. N.S., not significant; \*\* $P < 1.0 \times 10^{-2}$ ; \*\*\*\* $P < 1.0 \times 10^{-4}$  by random permutation experiments;  $n \geq 20$  independent larvae. (C) Representative changing patterns of head angles of larvae, taken every second for 1 min. Turning the head to left and right is presented as positive (+) and negative (−) angles, respectively. The patterns are obtained separately for larvae expressing each of the transgenes described in A. (D) Box plots showing the variability of head angle changes in larvae expressing the transgenes described in A. \*\*\*\* $P < 1.0 \times 10^{-4}$  by random permutation experiments;  $n = 4$  independent larvae.

protein toxicity in both dendrite morphology (Fig. 3 B and C) and larval locomotion.

**Coiled-Coil Structure-Mediated Target Sequestration Affinity of polyQ Proteins Strongly Correlates with polyQ-Induced Dendrite Defects.** Next, we further investigated the ability of Foxo to restore dendrite defects caused by MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins containing coiled-coil domains (see also Fig. 3 B and C). Foxo overexpression significantly ( $P < 0.05$ ) restored dendrite defects of C4da neurons expressing MJDtr-76Q, but the effect was weaker than that in C4da neurons expressing MJDtr-70Q<sub>cc0</sub> (Fig. 6 A and B), consistent with their efficacy in the restoration of larval locomotion defects. Foxo overexpression might affect the amounts of polyQ proteins. To test this possibility, we performed Western blot analysis of MJDtr-76Q and

MJDtr-70Q<sub>cc0</sub> with or without Foxo overexpression, and found no significant effect of Foxo overexpression on the amounts of both monomers and insoluble aggregates of the polyQ proteins (Fig. 6 C and D).

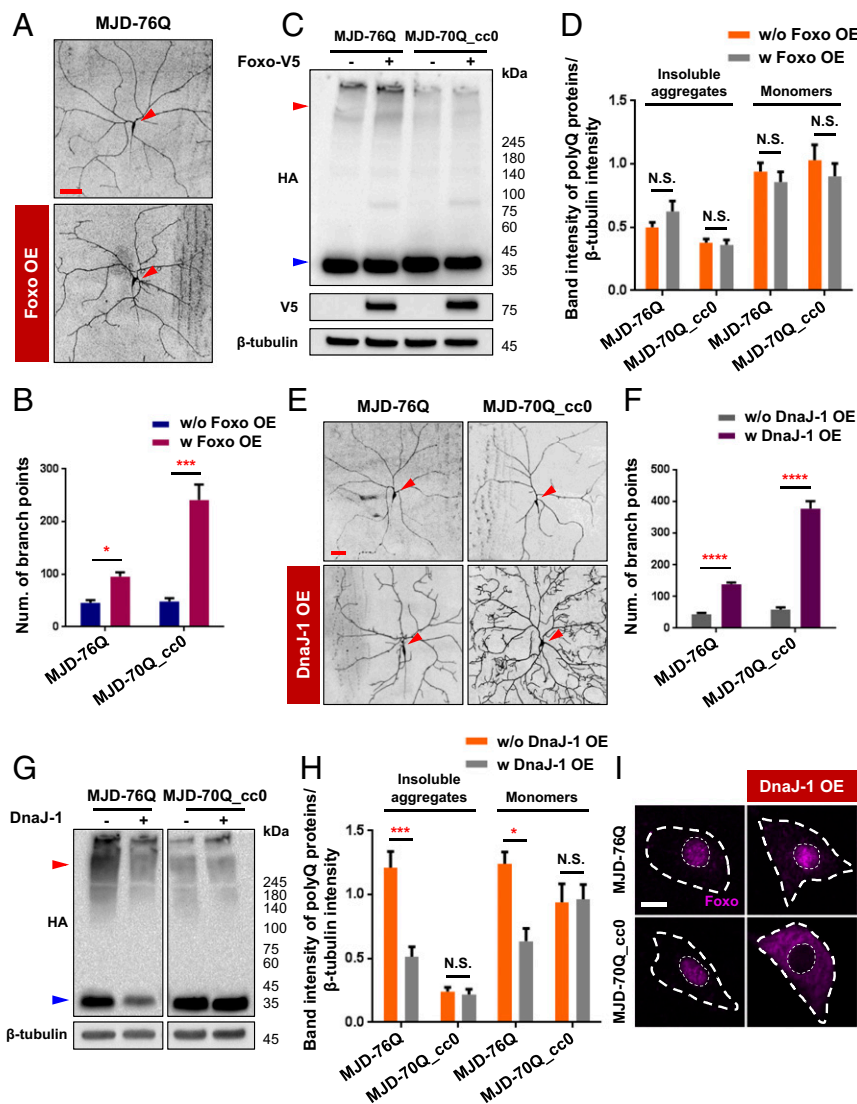
The difference in the restoration capability of Foxo between MJDtr-76Q- and MJDtr-70Q<sub>cc0</sub>-expressing C4da neurons might be due to the relative difference in polyQ toxicity. PolyQ toxicity is collectively determined by the amounts of toxic polyQ proteins and the affinity toward their sequestration targets. Previously, HSP-40, a mammalian homolog of DnaJ-1, was shown to affect both the amounts of toxic polyQ proteins and the interaction of polyQ proteins with their targets (29, 30). We first assessed the effect of DnaJ-1 overexpression on the extent of dendrite defects. Overexpression of DnaJ-1 in C4da neurons expressing MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> significantly ( $P < 1.0 \times 10^{-4}$ ) restored dendrite defects in both types of neurons (Fig. 6 E and F). However, this restoration was significantly stronger in C4da neurons expressing MJDtr-70Q<sub>cc0</sub> than in those expressing MJDtr-76Q, consistent with the restoration pattern observed after Foxo overexpression (Fig. 6B).

We then examined how DnaJ-1 overexpression altered the amounts of MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins. MJDtr-76Q proteins were significantly ( $P < 0.05$ ) reduced in the amounts of insoluble aggregates and monomers by DnaJ-1 overexpression (Fig. 6 G and H), consistent with the previous observation (29, 31). In contrast, MJDtr-70Q<sub>cc0</sub> proteins showed no significant changes. We next attempted to assess how DnaJ-1 overexpression affected the target sequestration affinity of MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins using co-IP. However, our co-IP experiments (Fig. 3F) showed virtually no interactions of Foxo with monomers of MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> proteins, making it difficult to accurately assess their target sequestration affinity. Thus, we instead measured Foxo subcellular localization to indirectly assess the target sequestration affinity of MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> proteins after co-overexpressing DnaJ-1 and Foxo in C4da neurons expressing MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> proteins. DnaJ-1 overexpression induced translocation of Foxo proteins from the nucleus to the cytoplasm in C4da neurons expressing MJDtr-70Q<sub>cc0</sub>, whereas only marginal changes in Foxo localization were observed in C4da neurons expressing MJDtr-76Q proteins (Fig. 6I).

In summary, DnaJ-1 overexpression resulted in a stronger decrease in the amount of MJDtr-76Q proteins, but a stronger decrease in target sequestration affinity of MJDtr-70Q<sub>cc0</sub> proteins. Considering a stronger restoration of dendrite defects by DnaJ-1 overexpression in C4da neurons expressing MJDtr-70Q<sub>cc0</sub> proteins than in those expressing MJDtr-76Q proteins, at least in the conditions tested in our study, these data suggest that the coiled-coil-mediated target sequestration affinity of toxic polyQ proteins appears to be more strongly associated with polyQ-induced dendrite defects, compared with the amount of toxic polyQ proteins.

## Discussion

In this study, among the coiled-coil-rich TFs implicated in dendrite morphogenesis, Foxo and Cut were selected as the most likely candidates to interact with polyQ proteins (Table 1). Of the two TFs, we focused on Foxo because we previously found that Cut indirectly interacted with MJDtr-78Q proteins through CBP (11). Nevertheless, another mode of interaction between Cut and SCA3 polyQ proteins based on coiled-coil structures could not be excluded. According to a previous study by Grueber et al. (20), the overexpression of Cut significantly increased dendrite branching in class I da (C1da) neurons. Based on this observation, we further examined dendrite branching in C1da neurons co-overexpressing Cut and MJDtr-76Q, MJDtr-70Q<sub>cc0</sub>, or MJDtr-70Q<sub>pQp</sub>. The overexpression of Cut alone elongated terminal dendrites and significantly ( $P < 1.0 \times 10^{-4}$ ) increased the



**Fig. 6.** Coiled-coil structure-mediated target sequestration affinity of polyQ proteins correlates with polyQ-induced dendrite defects. (A) Dendrite images of C4da neurons overexpressing MJDtr-76Q (Upper) or MJDtr-76Q + Foxo (Lower) (UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-76Q/+ and UAS-CD4td-GFP/UAS-Foxo;ppk-Gal4,UAS-MJDtr-76Q/+). Arrowheads (red) indicate cell bodies of C4da neurons. (Scale bar, 50  $\mu$ m.) (B) Quantification of the number of dendrite branch points in C4da neurons expressing the denoted transgenes (UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-76Q/+; UAS-CD4td-GFP/UAS-Foxo;ppk-Gal4,UAS-MJDtr-76Q/+; UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-70Q\_cc0/+; and UAS-CD4td-GFP/UAS-Foxo;ppk-Gal4,UAS-MJDtr-70Q\_cc0/+). The numbers of branch points are compared between C4da neurons expressing each SCA3 polyQ protein variant with and without Foxo overexpression. \* $P < 0.05$ ; \*\*\* $P < 1.0 \times 10^{-3}$  by two-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 5$  neurons. (C) Representative images obtained from Western blot analysis of adult fly heads (+/+;elav-GeneSwitch-Gal4/UAS-MJDtr-76Q, UAS-Foxo-V5/+;elav-GeneSwitch-Gal4/UAS-MJDtr-76Q, +/+;elav-GeneSwitch-Gal4/UAS-MJDtr-70Q\_cc0, and UAS-Foxo-V5/+; elav-GeneSwitch-Gal4/UAS-MJDtr-70Q\_cc0). SCA3 variants and Foxo-V5 were immunoblotted using anti-HA and anti-V5 antibodies, respectively.  $\beta$ -Tubulin is used as the loading control. Red and blue arrowheads indicate insoluble aggregates and monomers of SCA3 polyQ proteins, respectively.  $n = 3$  independent experiments. (D) Quantification of band intensities obtained from Western blot analysis (C) for insoluble aggregates and monomers of MJDtr-76Q and MJDtr-70Q\_cc0 proteins. Each band intensity was normalized with that of  $\beta$ -tubulin. N.S., not significant by two-way ANOVA with Tukey post hoc test; error bars, SEM;  $n = 3$  independent experiments. (E) Dendrite images of C4da neurons overexpressing MJDtr-76Q (Upper Left), MJDtr-70Q\_cc0 (Upper Right), MJDtr-76Q + DnaJ-1 (Lower Left), or MJDtr-70Q\_cc0 + DnaJ-1 (Lower Right) (UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-76Q/+; UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-70Q\_cc0/+; UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-70Q\_cc0/+; and UAS-CD4td-GFP/+;ppk-Gal4,UAS-MJDtr-70Q\_cc0/+). Arrowheads (red) indicate cell bodies of C4da neurons. (Scale bar, 50  $\mu$ m.) (F) Quantification of the number of dendrite branch points in C4da neurons expressing the transgenes described in D. \*\*\*\* $P < 1.0 \times 10^{-4}$  by two-way ANOVA with Tukey post hoc test; error bars, SEM;  $n \geq 3$  neurons. (G) Representative images obtained from Western blot analysis of adult fly heads (+/+;elav-Gal4/UAS-MJDtr-76Q, +/+;elav-Gal4,UAS-DnaJ-1/UAS-MJDtr-76Q, +/+;elav-Gal4/UAS-MJDtr-70Q\_cc0, and +/+;elav-Gal4,UAS-DnaJ-1/UAS-MJDtr-70Q\_cc0). SCA3 variants were immunoblotted using anti-HA antibody, and  $\beta$ -tubulin is used as the loading control. Red and blue arrowheads indicate insoluble aggregates and monomers of SCA3 polyQ proteins, respectively.  $n = 3$  independent experiments. (H) Quantification of band intensities obtained from Western blot analysis (G) for insoluble aggregates and monomers of MJDtr-76Q and MJDtr-70Q\_cc0 proteins. Each band intensity was normalized with that of  $\beta$ -tubulin. N.S., not significant; \* $P < 0.05$ ; \*\*\* $P < 1.0 \times 10^{-3}$  by two-way ANOVA with Tukey post hoc test; error bars, SEM;  $n = 3$  independent experiments. (I) Subcellular localization of Foxo (magenta) in C4da neurons overexpressing MJDtr-76Q (Upper Left), MJDtr-70Q\_cc0 (Lower Left), MJDtr-76Q + DnaJ-1 (Upper Right), or MJDtr-70Q\_cc0 + DnaJ-1 (Lower Right) (+/+;ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-76Q, UAS-DnaJ-1.K/+;ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-76Q, +/+;ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-70Q\_cc0, and UAS-DnaJ-1.K/+;ppk-Gal4,UASp-Foxo-GFP/UAS-MJDtr-70Q\_cc0). Outer and inner dashed lines (white) indicate the outlines of the cell body and nucleus, respectively. (Scale bar, 5  $\mu$ m.)



number of dendrite branch points (*SI Appendix, Fig. S6*), consistent with the previous findings (20). These changes in dendrites, however, were significantly ( $P < 1.0 \times 10^{-4}$ ) suppressed by co-overexpression of Cut with MJDtr-76Q or MJDtr-70Q<sub>cc0</sub>, but not with MJDtr-70Q<sub>pQp</sub> (*SI Appendix, Fig. S6B*). These results suggest that the function of Cut in dendrite branching is inhibited by SCA3 polyQ proteins containing coiled-coil structures, implying that such inhibition can be mediated by coiled-coil to coiled-coil interactions. Notably, CBP is also enriched with coiled-coil domains and thus likely to interact with polyQ proteins through the same interactions. Collectively, these data suggest that polyQ–CBP–Cut complex can be formed through coiled-coil to coiled-coil interactions between these proteins.

Our data showed that increased nuclear localization of Foxo is closely associated with polyQ protein toxicity (Fig. 3D). Foxo has been implicated in various neurodegenerative diseases, such as Alzheimer's (32, 33), prion (34), and amyotrophic lateral sclerosis diseases (35), suggesting potential associations with toxic proteins accumulated in these disorders. Foxo proteins have been observed in both the cytoplasm and nucleus in C4da neurons (36). In CCL39 fibroblast cells, Foxo proteins are predominantly localized in the cytoplasm, but upon inhibition of the PI3K–Akt signaling, they are dephosphorylated and translocated into the nucleus to induce the expression of their target genes (37). In our study, we observed that the co-overexpression of Foxo with MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> proteins increased the interaction of these proteins and the nuclear localization of Foxo (Fig. 3D), accompanied by reduced transcriptional activity of this TF (Fig. 3G and *SI Appendix, Fig. S3C*). The loss of Foxo function may indicate a reduced amount of the active TF—dephosphorylated Foxo—in the nucleus due to its interactions with MJDtr-76Q or MJDtr-70Q<sub>cc0</sub> proteins. Thus, our data suggest that MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins sequester Foxo in the nucleus, thereby reducing the amount of the active Foxo TF. Consistent with this model, Foxo RNAi led to larval locomotion defects (*SI Appendix, Fig. S5 B–E*). The overexpression of Foxo alone had no effect on larval locomotion (Fig. 5), suggesting that the amount of the active Foxo TF in the nucleus was not increased. Additional functional experiments should be carried out to test whether the overexpression of constitutively active Foxo TF contributes to larval locomotion.

We selected Foxo and Cut as the most likely TF targets of SCA3 polyQ proteins because they showed the strongest effects on dendrite branch points upon their knockdown or overexpression (Table 1). However, if the conventional statistical cut-off of  $P = 0.05$  from the analysis of variance (Tukey post hoc test) was applied, four additional TFs [Doublesex (Dsx), Nub, Relish (Rel), and Cnc] with statistically significant effects on dendrite branch points could also be selected on the basis of the knockdown and overexpression experiments (Fig. 3A and B). We did not pursue detailed functional experiments for these four TFs due to the lack of available antibodies or transgenic fly lines, and the difficulty in performing colocalization experiments because these TFs displayed the same nuclear localization as polyQ proteins. Nevertheless, the potential associations of these four TFs with neuronal processes impaired by protein toxicity have been previously reported. For example, Relish (NF- $\kappa$ B in mammals) was shown to regulate dendritic arborization (38), while the inhibition of NF- $\kappa$ B impaired synaptic signaling and learning in mice (39). Additionally, a number of studies have indicated the protective functions of Cnc (Nrf2 in mammals) against neuronal defects in various neurodegenerative diseases (40). Moreover, Dsx is involved in axonal projection in *Drosophila* (41), but its role in dendrite formation has been rarely studied. In addition, Nub is involved in the regulation of dendrite growth and branching (42). These data collectively suggest the potential roles of these coiled-coil-rich TFs in polyQ-induced

neuronal toxicity. Additional functional experiments are needed to elucidate their potential roles.

Comparison of dendrite defect restoration in C4da neurons expressing MJDtr-76Q and MJDtr-70Q<sub>cc0</sub> proteins suggests that the coiled-coil-mediated target sequestration affinity of toxic polyQ proteins is more strongly associated with polyQ-induced dendrite defects, compared with the amount of toxic polyQ proteins (Fig. 6). The severity of Foxo-dependent dendrite defects should be determined based on the amount of Foxo proteins sequestered by toxic polyQ proteins through coiled-coil to coiled-coil interactions (Fig. 4). The sequestered Foxo amount can be then determined by both the total amount (monomers and insoluble aggregates) and the coiled-coil-mediated target sequestration affinity of toxic polyQ proteins. First, regarding the total amount of toxic polyQ proteins, DnaJ-1 overexpression showed no effects on the amounts of monomers and insoluble aggregates in fly brains expressing MJDtr-70Q<sub>cc0</sub> containing only coiled-coil structures, but significant decreases in fly brains expressing MJDtr-76Q containing both coiled-coil and  $\beta$ -sheet structures. These data suggest that DnaJ-1 appears to predominantly affect the amount of toxic polyQ proteins influenced by  $\beta$ -sheet structures, although other biophysical properties of polyQ proteins may also contribute to the amount. This issue can be verified using a polyQ structural variant containing only  $\beta$ -sheet structures; however, to our knowledge, such a variant has not been developed due to the difficulty in breaking coiled-coil structures exclusively. Second, regarding the coiled-coil-mediated target sequestration affinity, DnaJ-1 overexpression showed only marginal effects on the target sequestration affinity in C4da neurons expressing MJDtr-76Q, but significantly decreases in C4da neurons expressing MJDtr-70Q<sub>cc0</sub>. These data suggest that DnaJ-1 appears to predominantly affect interactions of polyQ proteins with sequestered targets mediated by the coiled-coil structures.

Taking these data together, our study suggests that coiled-coil structures of polyQ proteins may serve as another fundamental structural basis, an alternative to  $\beta$ -sheet structures, for their interactions with targets containing coiled-coil domains, which can lead to polyQ pathology. Specifically, in this study, we propose the polyQ–Foxo interaction as a molecular mechanism for coiled-coil structure-mediated polyQ protein toxicity, such as dendrite and larval locomotion defects. Our study adds to the current understanding of the structure-based polyQ protein toxicity that mediates neuronal pathology through coiled-coil to coiled-coil interactions of these proteins with their TF targets. Furthermore, our findings suggest the potential of therapeutic approaches based on the inhibition of coiled-coil to coiled-coil interactions between pathogenic polyQ proteins and coiled-coil-rich TFs.

## Materials and Methods

**Fly stocks used were as follows:** *UAS-3xFLAG-Pros.S* (BL32245), *UASp-GFP-E2f1.WT* (BL34058), *UAS-Nej.WT-V5* (BL32573), *UAS-Flag-Rel.68* (BL55778), *UASp-Foxo-GFP* (BL43633), *UAS-Foxo.P* (BL9575), *UAS-Exex.B* (BL9929), *UAS-Pros RNAi* (BL42538), *UAS-Foxo RNAi* (BL32427), *UAS-Ct RNAi* (BL33967), *UAS-Kay RNAi* (BL33379), *UAS-Dsx RNAi* (BL55646), *UAS-Nub RNAi* (BL55305), *UAS-Nej RNAi* (BL37489), *UAS-Rel RNAi* (BL33661), *UAS-E2f1 RNAi* (BL36126), *UAS-Med RNAi* (BL31928), *UAS-Exex RNAi* (BL57709), *UAS-Cnc RNAi* (BL40854), *UAS-DnaJ-1.K* (BL30553), *UAS-MJDtr-27Q* (BL8149) and *elav-Gal4* (BL8760) were obtained from the Bloomington *Drosophila* Stock Center. *UAS-Cut-3xHA* (F001703), *UAS-Dsx-3xHA* (F001879), *UAS-Cnc-3xHA* (F000602), and *UAS-Med-3xHA* (F000591) were obtained from Fly ORF (43). The following lines were used as previously described: *UAS-mCD8-GFP* (III), *ppk-gal4* (III), *109(2)80-gal4* (II), *UAS-GMA*, *221-gal4* (III) (10), *UAS-Ct* (II), *ppk<sup>1a</sup>-gal4* (III), and *UAS-CD4-tdGFP* (II) (11). For pan-neuronal expression of transgenes, *elav-Gal4* was used. *109(2)80-gal4* was used to drive the expression of transgenes in da neuronal clusters. *ppk-gal4* and *ppk<sup>1a</sup>-gal4* were used for the expression of transgenes in C4da neurons. *221-gal4* was

used to expression of transgenes in C1da neurons. All flies were raised under standard conditions (25 °C and 60% humidity).

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